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NEWS 1 Web Page URLs for STN Seminar Schedule - N. America  
NEWS 2 Dec 17 The CA Lexicon available in the CAPLUS and CA files  
NEWS 3 Feb 06 Engineering Information Encompass files have new names  
NEWS 4 Feb 16 TOXLINE no longer being updated  
NEWS 5 Apr 23 Search Derwent WPINDEX by chemical structure  
NEWS 6 Apr 23 PRE-1967 REFERENCES NOW SEARCHABLE IN CAPLUS AND CA  
NEWS 7 May 07 DGENE Reload  
NEWS 8 Jun 20 Published patent applications (A1) are now in USPATFULL

NEWS EXPRESS May 23 CURRENT WINDOWS VERSION IS V6.0a,  
CURRENT MACINTOSH VERSION IS V5.0C (ENG) AND V5.0JB (JP),  
AND CURRENT DISCOVER FILE IS DATED 06 APRIL 2001

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FILE 'HOME' ENTERED AT 16:57:30 ON 01 JUL 2001

=> file biosis caplus embase medline cancerlit

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FILE 'MEDLINE' ENTERED AT 16:57:43 ON 01 JUL 2001

FILE 'CANCERLIT' ENTERED AT 16:57:43 ON 01 JUL 2001

=> s (ox? LDL)

4 FILES SEARCHED...  
L1 7898 (OX? LDL)

=> s l1 and antibod?

L2 1341 L1 AND ANTIBOD?

=> s l2 and fibrinogen

L3 22 L2 AND FIBRINOGEN

=> s l2 and plasminogen

L4 3L2 AND PLASMINOGEN

TI Method for detecting low density lipoprotein (LDL) or denatured LDL in blood  
 IN Uchida, Kazuo; Mashiba, Shinichi  
 PA Ikagaku Co., Ltd., Japan  
 SO Eur. Pat. Appl., 23 pp.  
 CODEN: EPXXDW  
 DT Patent  
 LA English  
 IC G01N033-92; C07K016-18  
 CC 9-10 (Biochemical Methods)  
 Section cross-reference(s): 14, 15

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	EP 1070962	A2	20010124	EP 2000-114984	20000720
	EP 1070962	A3	20010523		
	R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO				
	JP 2001091517	A2	20010406	JP 2000-12210	20000120
PRAI	JP 1999-207913	A	19990722		
	JP 2000-12210	A	20000120		
AB	A novel method for detecting LDL and denatured LDL (particularly, <b>oxidized LDL</b> ) having a significant concern with the onset and progression of arteriosclerosis and Alzheimer's disease is provided, wherein a complex of denatured LDL (particularly, <b>oxidized LDL</b> ) with an acute phase reactant, blood coagulation-fibrinolytic-related protein or disinfectant substance produced by macrophage is used as a measuring subject. Human LDL free of .alpha.1 antitrypsin and human fibronectin were treated with a copper sulfate soln. at 37.degree. over night to form an <b>oxidized LDL-fibronectin</b> complex. The complex was used as an immunogen in a mouse from which monoclonal <b>antibodies</b> were prepd. for use in assaying for the complex.				
ST	LDL lipoprotein detection blood; acute phase reactant denatured LDL detection; blood coagulation fibrinolytic related protein <b>oxidized LDL</b> detection; monoclonal <b>antibody oxidized LDL</b> fibronectin complex immunoassay				
IT	Apolipoproteins RL: BPR (Biological process); BIOL (Biological study); PROC (Process) (B, <b>antibody</b> to human; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)				
IT	Proteins, specific or class RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses) (C-reactive, complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)				
IT	Collagens, biological studies RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses) (LDL bonding to; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)				
IT	Diabetes mellitus (LDL- <b>fibrinogen</b> complex in; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)				
IT	Lipoproteins RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses) (Lp(a), complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)				
IT	Proteins, specific or class RL: ANT (Analyte); BPR (Biological process); SPN (Synthetic preparation); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses) (SAA (serum amyloid A), complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)				
IT	Glycoproteins, specific or class RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses) (SAP (serum amyloid, P), complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)				
IT	Proteins, specific or class RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)				

denatured LDL in blood)

IT Fibronectins  
 RL: ANT (Analyte); BPR (Biological process); SPN (Synthetic preparation);  
 THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study);  
 PREP (Preparation); PROC (Process); USES (Uses)  
 (complexes with LDL or denatured LDL; method for detecting low d.  
 lipoprotein (LDL) or denatured LDL in blood)

IT Complement  
**Fibrinogens**  
 Lactoferrins  
 .alpha.1-Acid glycoprotein  
 RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST  
 (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
 (complexes with LDL or denatured LDL; method for detecting low d.  
 lipoprotein (LDL) or denatured LDL in blood)

IT Enzymes, biological studies  
 RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical  
 study); BIOL (Biological study); USES (Uses)  
 (conjugates, with **antibodies**; method for detecting low d.  
 lipoprotein (LDL) or denatured LDL in blood)

IT Artery, disease  
 (coronary, **oxidized LDL** complexes in; method for  
 detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Macrophage  
 (disinfectant substance produced by, complexes with LDL or denatured  
 LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in  
 blood)

IT Immunoassay  
 (enzyme-linked immunosorbent assay; method for detecting low d.  
 lipoprotein (LDL) or denatured LDL in blood)

IT Immunoassay  
 (enzyme; method for detecting low d. lipoprotein (LDL) or denatured LDL  
 in blood)

IT Immunoassay  
 (immunoabsorption chromatog.; method for detecting low d. lipoprotein  
 (LDL) or denatured LDL in blood)

IT **Antibodies**  
 RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical  
 study); BIOL (Biological study); USES (Uses)  
 (labeled; method for detecting low d. lipoprotein (LDL) or denatured  
 LDL in blood)

IT Immunoassay  
 (latex agglutination test, latex flocculation; method for detecting low  
 d. lipoprotein (LDL) or denatured LDL in blood)

IT Lipoproteins  
 RL: ANT (Analyte); BPR (Biological process); SPN (Synthetic preparation);  
 THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study);  
 PREP (Preparation); PROC (Process); USES (Uses)  
 (low-d., complexes; method for detecting low d. lipoprotein (LDL) or  
 denatured LDL in blood)

IT Lipoproteins  
 RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST  
 (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
 (low-d., denatured; method for detecting low d. lipoprotein (LDL) or  
 denatured LDL in blood)

IT Lipoproteins  
 RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST  
 (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
 (low-d., oxidized; method for detecting low d. lipoprotein (LDL) or  
 denatured LDL in blood)

IT Lipoproteins  
 RL: ANT (Analyte); BPR (Biological process); RCT (Reactant); THU  
 (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC  
 (Process); USES (Uses)  
 (low-d.; method for detecting low d. lipoprotein (LDL) or denatured LDL  
 in blood)

IT Disinfectants  
 (macrophage-produced, complexes with LDL or denatured LDL; method for  
 detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Alzheimer's disease  
 Arteriosclerosis  
 Blood analysis

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IT Macrogloglobulins
RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST
  (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)
  (.alpha.2-, complexes with LDL or denatured LDL; method for detecting
  low d. lipoprotein (LDL) or denatured LDL in blood)
IT 9000-94-6D, Antithrombin, complexes with LDL or denatured LDL
  9001-26-7D, Prothrombin, complexes with LDL or denatured LDL 9001-63-2D,
  Lysozyme, complexes with LDL or denatured LDL 9001-91-6D,
  Plasminogen, complexes with LDL or denatured LDL 9002-04-4D,
  Thrombin, complexes with LDL or denatured LDL 9003-99-0D,
  Myeloperoxidase, complexes with LDL or denatured LDL 9035-58-9D,
  Blood-coagulation factor III, complexes with LDL or denatured LDL
  9041-92-3D, .alpha.1-Antitrypsin, complexes with LDL or denatured LDL
  140208-23-7D, Plasminogen activator inhibitor 1, complexes with
  LDL or denatured LDL 141176-92-3D, .alpha.1-Antichymotrypsin, complexes
  with LDL or denatured LDL
RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST
  (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)
  (method for detecting low d. lipoprotein (LDL) or denatured LDL in
  blood)

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=> d his

(FILE 'HOME' ENTERED AT 16:57:30 ON 01 JUL 2001)

FILE 'BIOSIS, CAPLUS, EMBASE, MEDLINE, CANCERLIT' ENTERED AT 16:57:43 ON  
01 JUL 2001

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L1 7898 S (OX? LDL)
L2 1341 S L1 AND ANTIBOD?
L3 22 S L2 AND FIBRINOGEN
L4 3 S L2 AND PLASMINOGEN
L5 4 S L2 AND LYSOZYME
L6 1 S L3 AND L4
L7 1 S L6 AND L5

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=> d l3 1-22 all

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L3 ANSWER 1 OF 22 BIOSIS COPYRIGHT 2001 BIOSIS
AN 2001:126543 BIOSIS
DN PREV200100126543
TI Circulating antibodies recognizing malondialdehyde-modified
  proteins in healthy subjects.
AU Vay, Daria; Parodi, Monica; Rolla, Roberta; Mottaran, Elisa; Vidali,
  Matteo; Bellomo, Giorgio; Albano, Emanuele (1)
CS (1) Department of Medical Science, University "Amedeo Avogadro" of East
  Piedmont, Via Solaroli 17, 28100, Novara: albano@med.unipmn.it Italy
SO Free Radical Biology & Medicine, (February 1, 2001) Vol. 30, No. 3, pp.
  277-286. print.
  ISSN: 0891-5849.
DT Article
LA English
SL English
AB Antibodies against malondialdehyde (MDA)-modified proteins are
  often increased in patients with diseases related to oxidative stress.
  However, the clinical significance of these antibodies is
  hampered by their frequent presence also in healthy controls. Aim of this
  work has been to characterize the immune reactivity against MDA-derived
  antigens in healthy subjects. The sera of 120 healthy subjects contained
  IgG and IgM targeting MDA-modified human albumin (HSA), fibrinogen
  , and LDL. These sera also displayed weak reactivity with oxidized
  LDL and HSA complexed with oxidized arachidonic acid. Conversely,
  oxidized HSA or HSA complexed with other aldehydic lipid peroxidation
  products was not recognized. Control sera also did not recognize cyclic
  dihydropyridine-MDA products, while HSA-MDA reactivity was associated ( $r > 0.9$ ;  $p < .0005$ ) with the presence of fluorescent lysine-conjugated-imine
  cross-links. In Western blots both IgG and IgM recognized high molecular
  weight HSA-MDA aggregates, but not monomeric HSA-MDA adducts. The addition
  of sodium cyanoborohydride, that prevented conjugated-imine fluorescence
  and protein aggregation during HSA-MDA preparation, abolished the
  antibody binding. This suggested that the plasma of healthy

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L3 ANSWER 2 OF 22 BIOSIS COPYRIGHT 2001 BIOSIS  
 AN 2000:182457 BIOSIS  
 DN PREV200000182457  
 TI Homocysteine and oxidized low density lipoprotein enhance platelet  
 adhesion to endothelial cells under flow conditions: Distinct mechanisms  
 of thrombogenic modulation.  
 AU Dardik, R. (1); Varon, D.; Tamarin, I.; Zivelin, A.; Salomon, O.;  
 Shenkman, B.; Savion, N.  
 CS (1) Sheba Medical Center, National Hemophilia Center and Institute of  
 Thrombosis and Hemostasis, Tel Hashomer, 52621 Israel  
 SO Thrombosis and Haemostasis, (Feb., 2000) Vol. 83, No. 2, pp. 338-344.  
 ISSN: 0340-6245.  
 DT Article  
 LA English  
 SL English  
 AB We investigated the effects of two well established risk factors for  
 cardiovascular disease, homocysteine and oxidized low density lipoprotein  
 (ox-LDL), on endothelial cell thrombogenicity. For  
 this purpose we studied platelet adhesion to human endothelial cells (EC)  
 under flow conditions at a shear rate of 350 s<sup>-1</sup> following EC treatment  
 with either homocysteine or ox-LDL. Treatment of EC  
 with either homocysteine (1 or 10 mmol/L for 16 h) or ox-  
 LDL (100 mug/ml for 16 h) resulted in a 2-3 fold enhancement in  
 platelet adhesion. The enhancement in platelet adhesion induced by 1  
 mmol/L homocysteine, but not that induced by 10 mmol/L homocysteine, was  
 absolutely dependent on fibrin formation. Homocysteine treatment has  
 significantly increased the cell surface tissue factor (TF) activity and  
 slightly reduced the expression of the intercellular adhesion molecule I  
 (ICAM-1). In contrast, ox-LDL treatment upregulated  
 ICAM-1 expression and had no significant effect on endothelial TF  
 activity. Neither homocysteine nor Ox-LDL affected  
 surface expression of the alphavbeta3 integrin. The homocysteine-induced  
 enhancement in platelet adhesion was almost completely abolished by  
 blockade of the EC TF activity by a polyclonal antibody. The  
 enhancing effect of homocysteine was also greatly reduced by inhibition of  
 the EC alphavbeta3 integrin, but was not affected by blockade of EC  
 ICAM-1. On the other hand, ox-LDL-induced enhancement  
 in platelet - EC adhesion was greatly inhibited by blocking ICAM-1 or  
 alphavbeta3, but remained unaffected by inhibition of TF activity.  
 Preincubation of platelets with the glycoprotein IIb-IIIa (GPIIb-IIIa)  
 antagonist Reo-Pro has virtually abolished the enhancing effect of both  
 homocysteine and ox-LDL. Our results suggest that  
 homocysteine and ox-LDL might increase endothelial  
 thrombogenicity by distinct mechanisms: homocysteine - by inducing TF  
 activity, and ox-LDL - by upregulating ICAM-1, both of  
 which enhance GPIIb-IIIa/fibrinogen dependent platelet adhesion  
 to EC. The alphavbeta3 integrin, although not affected by EC stimulation,  
 seems to play a crucial role in platelet-EC interaction regardless of the  
 mechanism of EC perturbation.  
 CC Cardiovascular System - General; Methods \*14501  
 Cytology and Cytochemistry - Human \*02508  
 Blood, Blood-Forming Organs and Body Fluids - General; Methods \*15001  
 Biochemical Studies - General \*10060  
 IT Major Concepts  
 Cardiovascular Medicine (Human Medicine, Medical Sciences); Hematology  
 (Human Medicine, Medical Sciences)  
 IT Parts, Structures, & Systems of Organisms  
 endothelial cells: thrombogenicity; platelet: blood and lymphatics  
 IT Chemicals & Biochemicals  
 alpha-V-beta-3 integrin; fibrin; fibrinogen; glycoprotein  
 IIb-IIIa; homocysteine; intercellular adhesion molecule-1; oxidized  
 low-density lipoprotein; tissue factor  
 IT Miscellaneous Descriptors  
 platelet adhesion  
 ORGN Super Taxa  
 Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia  
 ORGN Organism Name  
 human (Hominidae)  
 ORGN Organism Superterms  
 Animals; Chordates; Humans; Mammals; Primates; Vertebrates

SL . English

AB Although important roles of dietary n-3 fatty acids in the prevention of coronary heart disease (CHD) have been suggested, long-term effects of dietary alpha-linolenic acid (ALA, 18:3n-3) have not yet been established under controlled conditions. We tested whether a moderate increase of dietary ALA affects fatty acids composition in serum and the risk factors of CHD. **Oxidized LDL** (OxLDL) was directly measured by ELISA using **antibody** specific to OxLDL. By merely replacing soybean cooking oil (SO) with perilla oil (PO) (i.e., increasing 3 g/d of ALA), the n-6/n-3 ratio in the diet was changed from 4: 1 to 1 : 1. Twenty Japanese elderly subjects were initially given a SO diet for at least 6 mo (baseline period), a PO diet for 10 mo (intervention period), and then returned to the previous SO diet (washout period). ALA in the total serum lipid increased from 0.8 to 1.6% after 3 mo on the PO diet, but EPA and DHA increased in a later time, at 10 mo after the PO diet, from 2.5 to 3.6% and 5.3 to 6.4%, respectively (p < 0.05), and then returned to baseline in the washout period. In spite of increases of serum n-3 fatty acids, the OxLDL concentration did not change significantly when given the PO diet. Body weight, total serum cholesterol, triacylglycerol, glucose, insulin and HbA1c concentrations, platelet count and aggregation function, prothrombin time, partial thromboplastin time, **fibrinogen** and PAI-1 concentration, and other routine blood analysis did not change significantly when given the PO diet. These data indicate that, even in elderly subjects, a 3 g/d increase of dietary ALA could increase serum EPA and DHA in 10 mo without any major adverse effects.

CC Nutrition - General Studies, Nutritional Status and Methods \*13202

Physical Anthropology; Ethnobiology \*05000

Food Technology - General; Methods \*13502

Gerontology \*24500

Cardiovascular System - General; Methods \*14501

Biochemical Studies - General \*10060

IT Major Concepts

Nutrition

IT Diseases

coronary heart disease: heart disease

IT Chemicals & Biochemicals

alpha-linolenic acid: dietary intake; docosahexanoic acid: serum;

eicosapentanoic acid: serum; n-3 fatty acids: serum; **oxidized**

**LDL** [oxidized low density lipoprotein]

IT Alternate Indexing

Coronary Disease (MeSH)

IT Miscellaneous Descriptors

perilla oil: fats and oils

ORGN Super Taxa

Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia

ORGN Organism Name

human (Hominidae): Japanese, elderly

ORGN Organism Superterms

Animals; Chordates; Humans; Mammals; Primates; Vertebrates

RN 463-40-1 (ALPHA-LINOLENIC ACID)

L3 ANSWER 4 OF 22 BIOSIS COPYRIGHT 2001 BIOSIS

AN 1998:175330 BIOSIS

DN PREV199800175330

TI 3-nitrotyrosine in the proteins of human plasma determined by an ELISA method.

AU Khan, Jamshad; Brennan, David M.; Bradley, Nicholas; Gao, Beirong; Bruckdorfer, Richard; Jacobs, Michael (1)

CS (1) Dep. Pharmacol., Royal Free Hosp. Sch. Med., Rowland Hill St., London NW3 2PF UK

SO Biochemical Journal, (March 1, 1998) Vol. 330, No. 2, pp. 795-801.

ISSN: 0264-6021.

DT Article

LA English

AB The modification of tyrosine residues in proteins to 3-nitrotyrosine by peroxynitrite or other potential nitrating agents has been detected in biological systems that are subject to oxidative stress. A convenient semi-quantitative method has been developed to assay nitrated proteins in biological fluids and homogenates using a competitive ELISA developed in our laboratory. This assay selectivity detected 3-nitro-L-tyrosine residues in a variety of peroxynitrite-treated proteins (BSA, human serum albumin (HSA), alpha1-antiprotease inhibitor, pepsinogen and

CC Biochemical Studies - General \*10060  
 Biophysics - General Biophysical Studies \*10502  
 Enzymes - General and Comparative Studies; Coenzymes \*10802  
 Blood, Blood-Forming Organs and Body Fluids - General; Methods \*15001  
 Immunology and Immunochemistry - General; Methods \*34502  
 BC Hominidae 86215  
 IT Major Concepts  
     Biochemistry and Molecular Biophysics; Blood and Lymphatics (Transport and Circulation)  
 IT Parts, Structures, & Systems of Organisms  
     plasma: blood and lymphatics  
 IT Chemicals & Biochemicals  
     peroxynitrite; protein; BSA [bovine serum albumin]; LDL [low density lipoprotein]; 3-nitrotyrosine  
 IT Methods & Equipment  
     ELISA: determination method  
 ORGN Super Taxa  
     Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia  
 ORGN Organism Name  
     human (Hominidae)  
 ORGN Organism Superterms  
     Animals; Chordates; Humans; Mammals; Primates; Vertebrates  
 RN 3604-79-3 (3-NITROTYROSINE)  
 19059-14-4 (PEROXYNITRITE)

L3 ANSWER 5 OF 22 BIOSIS COPYRIGHT 2001 BIOSIS  
 AN 1997:105982 BIOSIS  
 DN PREV199799405185

TI Platelet integrin alpha-IIb-beta-3 (GPIIb-IIIa) is not implicated in the binding of LDL to intact resting platelets.  
 AU Pedreno, Javier (1); Fernandez, Rosa; Cullare, Cristina; Barcelo, Antonia; Elorza, Miguel Angel; De Castellarnau, Conxita  
 CS (1) Fundacio Invest. Cardiovasc., Hosp. Santa Creu Sant Pau, Avenida San Antonio Maria Claret 167, 08025 Barcelona Spain  
 SO Arteriosclerosis Thrombosis and Vascular Biology, (1997) Vol. 17, No. 1, pp. 156-163.  
 ISSN: 1079-5642.

DT Article  
 LA English

AB It has been suggested that the **fibrinogen** receptor (glycoprotein (GP) IIb-IIIa or platelet integrin alpha-IIb-beta-3) could be the binding site for low-density lipoprotein (LDL); however, recent data do not support this. Furthermore, GPIIb and not the GPIIb-IIIa complex is the main binding protein for lipoprotein(a) (Lp(a)). In the present study, we have investigated the interaction between Lp(a) particles and platelet LDL binding sites and whether platelet integrin alpha-IIb-beta-3 is implicated. Displacement experiments showed that 125I-LDL binding to intact resting platelets was inhibited with the same apparent affinity by both unlabeled LDL and apolipoprotein(a)-free lipoprotein particles (Lp(a)-, an LDL-like particle prepared from Lp(a)). Hill coefficients for displacement curves suggested that a single set of binding sites was involved. In contrast, both native and oxidized Lp(a) particles were unable to inhibit platelet LDL binding. Furthermore, platelets bound 125I-Lp(a)- particles to a class of saturable binding sites numbering approximately 1958+-235 binding sites per platelet with a dissociation constant (K-d) of 48.3+-12 times 10<sup>-9</sup> mol/L. These values were similar to those obtained for LDL. In contrast to Lp(a), evidence indicates that platelet integrin alpha-IIb-beta-3 was not involved in the interaction of LDL and intact resting platelets. First, specific ligands for platelet integrin alpha-IIb-beta-3, such as **fibrinogen**, vitronectin, and fibronectin, were unable to inhibit the binding of LDL to intact resting platelets. Second, similar LDL binding characteristics (K-d and B-max values) were found in platelets from control subjects and patients with type I and type II Glanzmann's thrombasthenia, characterized by total and partial lack of GPIIb-IIIa and **fibrinogen**, respectively. Third, polyclonal **antibodies** against the GPIIb-IIIa complex (edu-3 and 5B12), human antiserums against platelet alloantigens (anti-Bak-a/B and anti-PL-A1/2), anti-integrin subunits (anti-alpha-v and anti-beta-3), and a wide panel of monoclonal **antibodies** (mAbs) against well-known epitopes of GPIIb (M3, M4, M5, M6, and M95-2b) and GPIIIa (P23-7, P33, P37, P40, and P97) did not affect platelet LDL binding. Finally, in contrast to the proaggregatory effect of native and **oxidized**

Metabolism - Lipids \*13006  
 Metabolism - Proteins, Peptides and Amino Acids \*13012  
 Cardiovascular System - Physiology and Biochemistry \*14504  
 Cardiovascular System - Blood Vessel Pathology \*14508  
 Blood, Blood-Forming Organs and Body Fluids - Blood and Lymph Studies \*15002  
 Blood, Blood-Forming Organs and Body Fluids - Blood Cell Studies \*15004  
 Blood, Blood-Forming Organs and Body Fluids - Blood, Lymphatic and Reticuloendothelial Pathologies \*15006  
 BC Hominidae \*86215  
 IT Major Concepts  
   Biochemistry and Molecular Biophysics; Bioenergetics (Biochemistry and Molecular Biophysics); Blood and Lymphatics (Transport and Circulation); Cardiovascular Medicine (Human Medicine, Medical Sciences); Cardiovascular System (Transport and Circulation); Cell Biology; Hematology (Human Medicine, Medical Sciences); Membranes (Cell Biology); Metabolism  
 IT Chemicals & Biochemicals  
   INTEGRIN  
 IT Miscellaneous Descriptors  
   ATHEROSCLEROSIS; BINDING; BIOCHEMISTRY AND BIOPHYSICS; BLOOD AND LYMPHATICS; **FIBRINOGEN** RECEPTOR; INTACT RESTING PLATELETS; LDL; LDL BINDING SITES; LIPOPROTEIN; LIPOPROTEIN(A); LOW-DENSITY LIPOPROTEIN; LOW-DENSITY LIPOPROTEIN BINDING SITES; OXIDATION; OXIDIZED LIPOPROTEINS; PLATELET GLYCOPROTEIN IIB-IIIA; PLATELET GPIIB-IIIA; PLATELET INTEGRIN-ALPHA-IIB,BETA-3; VASCULAR DISEASE  
 ORGN Super Taxa  
   Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia  
 ORGN Organism Name  
   human (Hominidae)  
 ORGN Organism Superterms  
   animals; chordates; humans; mammals; primates; vertebrates  
 RN 153-87-7Q (INTEGRIN)  
   60791-49-3Q (INTEGRIN)

L3 ANSWER 6 OF 22 CAPLUS COPYRIGHT 2001 ACS  
 AN 2001:93225 CAPLUS  
 DN 134:264876

TI Circulating **antibodies** recognizing malondialdehyde-modified proteins in healthy subjects

AU Vay, D.; Parodi, M.; Rolla, R.; Mottaran, E.; Vidali, M.; Bellomo, G.; Albano, E.

CS Department of Medical Sciences, University "Amedeo Avogadro" of East Piedmont, Novara, Italy

SO Free Radical Biol. Med. (2001), 30(3), 277-286  
 CODEN: FRBMEH; ISSN: 0891-5849

PB Elsevier Science Inc.

DT Journal

LA English

CC 15-3 (Immunochemistry)

AB **Antibodies** against malondialdehyde (MDA)-modified proteins are often increased in patients with diseases related to oxidative stress. However, the clin. significance of these **antibodies** is hampered by their frequent presence also in healthy controls. Aim of this work has been to characterize the immune reactivity against MDA-derived antigens in healthy subjects. The sera of 120 healthy subjects contained IgG and IgM targeting MDA-modified human albumin (HSA), **fibrinogen**, and LDL. These sera also displayed weak reactivity with **oxidized LDL** and HSA complexed with oxidized arachidonic acid. Conversely, oxidized HSA or HSA complexed with other aldehydic lipid peroxidn. products was not recognized. Control sera also did not recognize cyclic dihydropyridine-MDA products, while HSA-MDA reactivity was assocd. ( $r > 0.9$ ;  $p < .0005$ ) with the presence of fluorescent lysine-conjugated-imine cross-links. In Western blots both IgG and IgM recognized high mol. wt. HSA-MDA aggregates, but not monomeric HSA-MDA adducts. The addn. of sodium cyanoborohydride, that prevented conjugated-imine fluorescence and protein aggregation during HSA-MDA prepn., abolished the **antibody** binding. This suggested that the plasma of healthy subjects contained IgG and IgM recognizing protein aggregates linked through 1-amino-3-imino-propene bridges. The function of these **antibodies** is at the moment unknown, but they might contribute to scavenging MDA cross-linked proteins.



(low-d., malondialdehyde-modified; circulating **antibodies** recognizing malondialdehyde-modified proteins in healthy humans)

IT Lipoproteins  
 RL: BPR (Biological process); BIOL (Biological study); PROC (Process)  
 (low-d., oxidized, malondialdehyde-modified; circulating **antibodies** recognizing malondialdehyde-modified proteins in healthy humans)

IT Albumins, biological studies  
**Fibrinogens**  
 Proteins, specific or class  
 RL: BPR (Biological process); BIOL (Biological study); PROC (Process)  
 (malondialdehyde-modified; circulating **antibodies** recognizing malondialdehyde-modified proteins in healthy humans)

IT Lipids, biological studies  
 RL: BPR (Biological process); BIOL (Biological study); PROC (Process)  
 (peroxidn.; circulating **antibodies** recognizing malondialdehyde-modified proteins in healthy humans in relation to)

IT 506-32-1D, Arachidonic acid, oxidized  
 RL: BPR (Biological process); BIOL (Biological study); PROC (Process)  
 (malondialdehyde-modified; circulating **antibodies** recognizing malondialdehyde-modified proteins in healthy humans)

RE.CNT 38

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L3 ANSWER 7 OF 22 CAPLUS COPYRIGHT 2001 ACS

AN 2001:62437 CAPLUS

DN 134:97520

TI Method for detecting low density lipoprotein (LDL) or denatured LDL in blood

IN Uchida, Kazuo; Mashiba, Shinichi

PA Ikagaku Co., Ltd., Japan

SO Eur. Pat. Appl., 23 pp.

CODEN: EPXXDW

DT Patent

LA English

oxidized LDL) with an acute phase reactant, blood coagulation-fibrinolytic-related protein or disinfectant substance produced by macrophage is used as a measuring subject. Human LDL free of .alpha.1 antitrypsin and human fibronectin were treated with a copper sulfate soln. at 37.degree. over night to form an **oxidized LDL-fibronectin complex**. The complex was used as an immunogen in a mouse from which monoclonal **antibodies** were prepd. for use in assaying for the complex.

ST LDL lipoprotein detection blood; acute phase reactant denatured LDL detection; blood coagulation fibrinolytic related protein **oxidized LDL** detection; monoclonal **antibody oxidized LDL** fibronectin complex immunoassay

IT Apolipoproteins  
 RL: BPR (Biological process); BIOL (Biological study); PROC (Process)  
 (B, **antibody** to human; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Proteins, specific or class  
 RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
 (C-reactive, complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Collagens, biological studies  
 RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)  
 (LDL bonding to; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Diabetes mellitus  
 (LDL-**fibrinogen** complex in; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Lipoproteins  
 RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
 (Lp(a), complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Proteins, specific or class  
 RL: ANT (Analyte); BPR (Biological process); SPN (Synthetic preparation); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)  
 (SAA (serum amyloid A), complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Glycoproteins, specific or class  
 RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
 (SAP (serum amyloid, P), complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Proteins, specific or class  
 RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
 (acute-phase, complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Lipoproteins  
 RL: ANT (Analyte); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)  
 (arteriosclerosis-assocd.; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Proteins, specific or class  
 RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
 (basic, complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Proteins, specific or class  
 RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
 (blood coagulation-fibrinolytic-related, complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Fibronectins  
 RL: ANT (Analyte); BPR (Biological process); SPN (Synthetic preparation); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)  
 (complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Complement

LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Immunoassay  
(enzyme-linked immunosorbent assay; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Immunoassay  
(enzyme; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Immunoassay  
(immunoabsorption chromatog.; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT **Antibodies**  
RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)  
(labeled; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Immunoassay  
(latex agglutination test, latex flocculation; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Lipoproteins  
RL: ANT (Analyte); BPR (Biological process); SPN (Synthetic preparation); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)  
(low-d., complexes; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Lipoproteins  
RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
(low-d., denatured; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Lipoproteins  
RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
(low-d., oxidized; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Lipoproteins  
RL: ANT (Analyte); BPR (Biological process); RCT (Reactant); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
(low-d.; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Disinfectants  
(macrophage-produced, complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Alzheimer's disease  
Arteriosclerosis  
Blood analysis  
Hybridoma  
Immunoassay  
(method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT **Antibodies**  
RL: ARG (Analytical reagent use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)  
(method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT **Antibodies**  
RL: ARG (Analytical reagent use); BPN (Biosynthetic preparation); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PREP (Preparation); PROC (Process); USES (Uses)  
(monoclonal, to LDL-fibronectin complex; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Disease, animal  
(syndrome X, LDL-fibrinogen complex in; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT Macroglobulins  
RL: ANT (Analyte); BPR (Biological process); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); PROC (Process); USES (Uses)  
(.alpha.2-, complexes with LDL or denatured LDL; method for detecting low d. lipoprotein (LDL) or denatured LDL in blood)

IT 9000-94-6D, Antithrombin, complexes with LDL or denatured LDL  
9001-26-7D, Prothrombin, complexes with LDL or denatured LDL 9001-63-2D, Lysozyme, complexes with LDL or denatured LDL 9001-91-6D, Plasminogen,

of thrombogenic modulation

AU Dardik, R.; Varon, D.; Tamarin, I.; Zivelin, A.; Salomon, O.; Shenkman, B.; Savion, N.

CS National Hemophilia Center, Sheba Medical Center, Tel Hashomer, 52621, Israel

SO Thromb. Haemostasis (2000), 83(2), 338-344  
CODEN: THHADQ; ISSN: 0340-6245

PB F. K. Schattauer Verlagsgesellschaft mbH

DT Journal

LA English

CC 14-5 (Mammalian Pathological Biochemistry)

AB The authors investigated the effects of 2 the authors 11 established risk factors for cardiovascular disease, homocysteine and oxidized low d. lipoprotein (**ox-LDL**), on endothelial cell thrombogenicity. For this purpose the authors studied platelet adhesion to human endothelial cells (EC) under flow conditions at a shear rate of 350 s<sup>-1</sup> following EC treatment with either homocysteine or **ox-LDL**. Treatment of EC with either homocysteine (1 or 10 mmol/L for 16 h) or **ox-LDL** (100 .mu.g/mL for 16 h) resulted in a 2-3-fold enhancement in platelet adhesion. The enhancement in platelet adhesion induced by 1 mmol/L homocysteine, but not that induced by 10 mmol/L homocysteine, was absolutely dependent on fibrin formation. Homocysteine treatment has significantly increased the cell surface tissue factor (TF) activity and slightly reduced the expression of the intercellular adhesion mol. 1 (ICAM-1). In contrast, **ox-LDL** treatment upregulated ICAM-1 expression and had no significant effect on endothelial TF activity. Neither homocysteine nor **ox-LDL** affected surface expression of the .alpha.v.beta.3 integrin. The homocysteine-induced enhancement in platelet adhesion was almost completely abolished by blockade of the EC TF activity by a polyclonal **antibody**. The enhancing effect of homocysteine was also greatly reduced by inhibition of the EC .alpha.v.beta.3 integrin, but was not affected by blockade of EC ICAM-1. On the other hand, **ox-LDL**-induced enhancement in platelet-EC adhesion was greatly inhibited by blocking ICAM-1 or .alpha.v.beta.3, but remained unaffected by inhibition of TF activity. Preincubation of platelets with the glycoprotein IIb-IIIa (GPIIb-IIIa) antagonist Reo-Pro has virtually abolished the enhancing effect of both homocysteine and **ox-LDL**. These results suggest that homocysteine and **ox-LDL** might increase endothelial thrombogenicity by distinct mechanisms: homocysteine - by inducing TF activity, and **ox-LDL** - by upregulating ICAM-1, both of which enhance GPIIb-IIIa/**fibrinogen** dependent platelet adhesion to EC. The .alpha.v.beta.3 integrin, although not affected by EC stimulation, seems to play a crucial role in platelet-EC interaction regardless of the mechanism of EC perturbation.

ST homocysteine oxLDL platelet adhesion endothelium thrombosis; low density lipoprotein oxidized ICAM1 platelet adhesion endothelium thrombosis; tissue factor homocysteine platelet adhesion endothelium thrombosis; GPIIbIIIa **fibrinogen** platelet adhesion endothelium thrombosis  
homocysteine oxLDL

IT Cell adhesion molecules  
RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BIOL (Biological study); PROC (Process)  
(ICAM-1 (intercellular adhesion mol. 1); **ox-LDL** increased endothelial thrombogenicity via ICAM-1 enhancing GPIIb-IIIa/**fibrinogen** dependent platelet adhesion)

IT Platelet (blood)  
(adhesion; homocysteine and **ox-LDL** increased endothelial thrombogenicity via TF and ICAM-1 enhancing GPIIb-IIIa/**fibrinogen** dependent platelet adhesion)

IT Blood vessel  
(endothelium; homocysteine and **ox-LDL** increased endothelial thrombogenicity via TF and ICAM-1 enhancing GPIIb-IIIa/**fibrinogen** dependent platelet adhesion)

IT Thrombosis  
(homocysteine and **ox-LDL** increased endothelial thrombogenicity via TF and ICAM-1 enhancing GPIIb-IIIa/**fibrinogen** dependent platelet adhesion)

IT **Fibrinogens**  
RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BIOL (Biological study); PROC (Process)

(homocysteine and **ox-LDL** increased endothelial thrombogenicity via TF enhancing GPIIb-IIIa/**fibrinogen** dependent platelet adhesion)

IT 9035-58-9, Blood-coagulation factor III  
RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BIOL (Biological study); PROC (Process)  
(homocysteine increased endothelial thrombogenicity via TF enhancing GPIIb-IIIa/**fibrinogen** dependent platelet adhesion)

RE.CNT 30

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L3 ANSWER 9 OF 22 CAPLUS COPYRIGHT 2001 ACS

AN 1998:226448 CAPLUS

DN 128:305846

TI 3-Nitrotyrosine in the proteins of human plasma determined by an ELISA method

AU Khan, Jamshad; Brennand, David M.; Bradley, Nicholas; Gao, Beirong; Bruckdorfer, Richard; Jacobs, Michael

CS Department of Pharmacology, Royal Free Hospital School of Medicine, London, NW3 2PF, UK

SO Biochem. J. (1998), 330(2), 795-801

CODEN: BIJOAK; ISSN: 0264-6021

PB Portland Press Ltd.

DT Journal

LA English

CC 9-10 (Biochemical Methods)

AB The modification of tyrosine residues in proteins to 3-nitrotyrosine by peroxynitrite or other potential nitrating agents has been detected in biol. systems that are subject to oxidative stress. A convenient semi-quant. method has been developed to assay nitrated proteins in biol. fluids and homogenates using a competitive ELISA developed in our lab. This assay selectivity detected 3-nitro-L-tyrosine residues in a variety of peroxynitrite-treated proteins (BSA, human serum albumin (HSA), .alpha.1-antiprotease inhibitor, pepsinogen and **fibrinogen**) and also in a nitrated peptide, but had a low affinity for free 3-nitro-L-tyrosine and 3-chloro-L-tyrosine. The IC50 values for the inhibition of **antibody** binding by different nitrated proteins were in the range 5-100 nM, suggesting that the **antibody** discriminated between nitrotyrosine residues in different environments. The presence of nitrotyrosine in plasma proteins was detected by Western blot anal and quantified by the ELISA. A concn. of 0.12+-0.01 .mu.M nitro-BSA equiv. was measured in the proteins of normal plasma which was increased in peroxynitrite-treated plasma and was elevated in inflammatory

conditions. HSA and low-d. lipoprotein (LDL) isolated from plasma contained  $0.085 \pm 0.04$  and  $0.03 \pm 0.006$  nmol nitro-BSA equiv./mg protein, resp. Comparison of the level of nitration in peroxynitrite-treated HSA and LDL in the presence and absence of plasma indicates that nitration and presumably oxidn. is inhibited by plasma antioxidants. The presence of nitrotyrosine in LDL is consistent with previous reports implicating peroxynitrite in the oxidative modification of lipoproteins and the presence of a low concn. of **oxidized LDL** in the blood.

ST nitrotyrosine protein plasma detd ELISA  
IT Blood analysis

RL: ANT (Analyte); ANST (Analytical study)  
(3-Nitrotyrosine in proteins of human plasma detd. by ELISA method)

L3 ANSWER 10 OF 22 CAPLUS COPYRIGHT 2001 ACS

AN 1997:115499 CAPLUS

DN 126:142448

TI Platelet integrin .alpha.IIb.beta.3 (GPIIb-IIIa) is not implicated in the binding of LDL to intact resting platelets

AU Pedreno, Javier; Fernandez, Rosa; Cullare, Cristina; Barcelo, Antonia; Elorza, Miguel Angel; De Castellarnau, Conxita

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SO Arterioscler., Thromb., Vasc. Biol. (1997), 17(1), 156-163

CODEN: ATVBFA; ISSN: 1079-5642

PB American Heart Association

DT Journal

LA English

CC 13-5 (Mammalian Biochemistry)

AB It has been suggested that the **fibrinogen** receptor (glycoprotein IIb-IIIa or platelet integrin .alpha.IIb.beta.3) could be the binding site for low-d. lipoprotein (LDL); however, recent data do not support this. Furthermore, GPIIb and not the GPIIb-IIIa complex is the main binding protein for lipoprotein(a) [Lp(a)]. In the present study, the interaction between Lp(a) particles and platelet LDL binding sites and the role of platelet integrin .alpha.IIb.beta.3 were investigated. Displacement expts. showed that 125I-LDL binding to intact resting platelets was inhibited with the same apparent affinity by both unlabeled LDL and apolipoprotein(a)-free lipoprotein particles Lp(a)- [an LDL-like particle prepd. from Lp(a)]. Hill coeffs. for displacement curves suggested that a single set of binding sites was involved. In contrast, both native and oxidized Lp(a) particles were unable to inhibit platelet LDL binding. Furthermore, platelets bound 125I-Lp(a)- particles to a class of saturable binding sites numbering approx. 1958.+-.235 binding sites per platelet with a disocn. const. (Kd) of 48.3.+-.12.times.10<sup>-9</sup> mol/L. These values were similar to those obtained for LDL. In contrast to Lp(a), evidence indicates that platelet integrin .alpha.IIb.beta.3 was not involved in the interaction of LDL and intact resting platelets. First, specific ligands for platelet integrin .alpha.IIb.beta.3, such as **fibrinogen**, vitronectin, and fibronectin, were unable to inhibit the binding of LDL to intact resting platelets. Second, similar LDL binding characteristics (Kd and Bmax values) were found in platelets from control subjects and patients with type I and type II Glanzmann's thrombasthenia, characterized by total and partial lack of GPIIb-IIIa and **fibrinogen**, resp. Third, polyclonal **antibodies** against the GPIIb-IIIa complex (edu-3 and 5B12), human antisera against platelet alloantigens (anti-Baka/B and anti-PLA1/2), anti-integrin subunits (anti-.alpha.v and anti-.beta.3), and a wide panel of monoclonal **antibodies** against well-known epitopes of GPIIb (M3, M4, M5, M6, and M95-2b) and GPIIIa (P23-7, P33, P37, P40, and P97) did not affect platelet LDL binding. Finally, in contrast to the proaggregatory effect of native and **oxidized LDL**, both native and oxidized Lp(a) particles caused a significant dose-dependent decrease of collagen-induced platelet aggregation. In conclusion, It is demonstrated that neither the GPIIb-IIIa complex nor GPIIb and GPIIIa individually are membrane binding proteins for LDL on intact resting platelets. Lp(a) particles do not interact with platelet LDL binding sites, and their biol. response is clearly different from that of LDL.

ST platelet LDL binding integrin glycoprotein

IT Glycoproteins (specific proteins and subclasses)

RL: BSU (Biological study, unclassified); BIOL (Biological study)  
(IIb/IIIa; platelet integrin .alpha.IIb.beta.3 (GPIIb-IIIa) is not implicated in binding of LDL to intact resting platelets)

IT Platelet (blood)

(platelet integrin .alpha.IIb.beta.3 (GPIIb-IIIa) is not implicated in binding of LDL to intact resting platelets)

IT Low-density lipoproteins

RL: BPR (Biological process); BIOL (Biological study); PROC (Process)  
(platelet integrin .alpha.IIb.beta.3 (GPIIb-IIIa) is not implicated in binding of LDL to intact resting platelets)

IT Integrin .alpha.IIb.beta.3

RL: BSU (Biological study, unclassified); BIOL (Biological study)  
(platelet integrin .alpha.IIb.beta.3 (GPIIb-IIIa) is not implicated in

FS 006 Internal Medicine  
018 Cardiovascular Diseases and Cardiovascular Surgery  
030 Pharmacology  
037 Drug Literature Index

LA English

SL English

AB We measured and compared levels of platelet-derived microparticles (PMPs), monocyte-derived microparticles (MMPs), CD62P on activated platelets, soluble E-selectin (sE-selectin), and antioxidized low density lipoprotein (LDL) **antibody** in hyperlipidemia patients and control subjects. Binding of anti-GPIIb/IIIa and anti-GPIb monoclonal **antibodies** to platelets was not significantly different between hyperlipidemia patients and controls. However, expression of CD62P on platelets and levels of PMPs were higher for hyperlipidemia patients than in controls, although the difference between groups in CD62P expression was not significant (PMPs: 534  $\pm$  63 vs. 388  $\pm$  47,  $p < 0.05$ ; CD62P: 9.1%  $\pm$  1.45 vs. 7.3%  $\pm$  1.15, N.S.). Although there were no differences in expression of CD36 and CD40 by monocytes between the two groups, levels of MMPs were higher in hyperlipidemia patients than in controls (MMPs: 147  $\pm$  21 vs. 59  $\pm$  8, respectively,  $p < 0.01$ ). Levels of anti-**oxidized LDL antibody** and sE-selectin were also higher in hyperlipidemia patients. We studied the effects of Saiko-ka-ryukotsu-borei-to on levels of these factors in patients with elevated triglyceride levels. After Saiko-ka-ryukotsu-borei-to treatment, levels of CD62P, PMPs, sE-selectin, and anti-**oxidized LDL antibody** were reduced significantly. Levels of triglycerides, total cholesterol and MMPs also decreased, but the changes were not significant. These findings suggest that Saiko-ka-ryukotsu-borei-to prevents the development of vascular complications in hyperlipidemia patients.

CT Medical Descriptors:

\*hyperlipidemia: DT, drug therapy  
drug effect

measurement

antigen binding

protein expression

triacylglycerol blood level

cholesterol blood level

vascular disease: CO, complication

vascular disease: DT, drug therapy

vascular disease: PC, prevention

monocyte

thrombocyte activation

human

male

female

clinical article

controlled study

aged

adult

article

priority journal

Drug Descriptors:

\*saiko ka ryukotsu borei to: DT, drug therapy

\*saiko ka ryukotsu borei to: PD, pharmacology

endothelial leukocyte adhesion molecule 1

**monoclonal antibody**

**fibrinogen receptor: EC, endogenous compound**

PADGEM protein: EC, endogenous compound

oxidized low density lipoprotein: EC, endogenous compound

**protein antibody: EC, endogenous compound**

**oxidized low density lipoprotein antibody: EC, endogenous compound**

CD36 antigen: EC, endogenous compound

CD40 antigen: EC, endogenous compound

triacylglycerol: EC, endogenous compound

cholesterol: EC, endogenous compound

glycoprotein Ib: EC, endogenous compound

unclassified drug

RN (endothelial leukocyte adhesion molecule 1) 128875-25-2; (cholesterol)  
57-88-5

L3 ANSWER 12 OF 22 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.



However, the clinical significance of these **antibodies** is hampered by their frequent presence also in healthy controls. Aim of this work has been to characterize the immune reactivity against MDA-derived antigens in healthy subjects. The sera of 120 healthy subjects contained IgG and IgM targeting MDA-modified human albumin (HSA), **fibrinogen**, and LDL. These sera also displayed weak reactivity with **oxidized LDL** and HSA complexed with oxidized arachidonic acid. Conversely, oxidized HSA or HSA complexed with other aldehydic lipid peroxidation products was not recognized. Control sera also did not recognize cyclic dihydropyridine-MDA products, while HSA-MDA reactivity was associated ( $r > 0.9$ ;  $p < .0005$ ) with the presence of fluorescent lysine-conjugated-imine cross-links. In Western blots both IgG and IgM recognized high molecular weight HSA-MDA aggregates, but not monomeric HSA-MDA adducts. The addition of sodium cyanoborohydride, that prevented conjugated-imine fluorescence and protein aggregation during HSA-MDA preparation, abolished the **antibody** binding. This suggested that the plasma of healthy subjects contained IgG and IgM recognizing protein aggregates linked through 1-amino-3-imino-propene bridges. The function of these **antibodies** is at the moment unknown, but they might contribute to scavenging MDA cross-linked proteins. .COPYRGT. 2001 Elsevier Science Inc.

CT

Medical Descriptors:

\*antigen recognition  
oxidative stress  
immunoreactivity  
lipid peroxidation  
molecular weight  
antigen binding  
Western blotting  
protein aggregation  
oxidation  
correlation function  
antigenicity  
concentration response  
adsorption  
cross linking  
protein modification  
human  
male  
female  
normal human  
controlled study  
adult  
article  
priority journal

Drug Descriptors:

\*malonaldehyde  
**\*protein antibody: EC, endogenous compound**  
\*malonaldehyde modified protein: EC, endogenous compound  
**\*malonaldehyde modified protein antibody: EC, endogenous compound**  
\*protein: EC, endogenous compound  
immunoglobulin A: EC, endogenous compound  
immunoglobulin M: EC, endogenous compound  
immunoglobulin G: EC, endogenous compound  
human serum albumin  
**fibrinogen**  
low density lipoprotein  
arachidonic acid  
cyanoborohydride sodium  
reducing agent  
apolipoprotein B  
free radical  
antigen: EC, endogenous compound  
cyclic dihydropyridine: EC, endogenous compound  
unclassified drug

RN

(malonaldehyde) 542-78-9; (protein) 67254-75-5; (immunoglobulin M) 9007-85-6; (immunoglobulin G) 97794-27-9; (human serum albumin) 9048-49-1; (**fibrinogen**) 9001-32-5; (arachidonic acid) 506-32-1, 6610-25-9, 7771-44-0; (cyanoborohydride sodium) 25895-60-7, 33195-00-5

L3

ANSWER 13 OF 22 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN

2000095337 EMBASE

TI

Long-term effects of dietary .alpha.-linolenic acid from perilla oil on

dietary .alpha.-linolenic acid (ALA, 18:3n-3) have not yet been established under controlled conditions. We tested whether a moderate increase of dietary ALA affects fatty acids composition in serum and the risk factors of CHD. Oxidized LDL (OxLDL) was directly measured by ELISA using antibody specific to OxLDL. By merely replacing soybean cooking oil (SO) with perilla oil (PO) (i.e., increasing 3 g/d of ALA), the n-6/n-3 ratio in the diet was changed from 4:1 to 1:1. Twenty Japanese elderly subjects were initially given a SO diet for at least 6 mo (baseline period), a PO diet for 10 mo (intervention period), and then returned to the previous SO diet (washout period). ALA in the total serum lipid increased from 0.8 to 1.6% after 3 mo on the PO diet, but EPA and DHA increased in a later time, at 10 mo after the PO diet, from 2.5 to 3.6% and 5.3 to 6.4%, respectively ( $p < 0.05$ ), and then returned to baseline in the washout period. In spite of increases of serum n-3 fatty acids, the OxLDL concentration did not change significantly when given the PO diet. Body weight, total serum cholesterol, triacylglycerol, glucose, insulin and HbA1c concentrations, platelet count and aggregation function, prothrombin time, partial thromboplastin time, **fibrinogen** and PAI-1 concentration, and other routine blood analysis did not change significantly when given the PO diet. These data indicate that, even in elderly subjects, a 3 g/d increase of dietary ALA could increase serum EPA and DHA in 10 mo without any major adverse effects.

CT Medical Descriptors:

\*lipid composition  
 \*dietary intake  
 \*ischemic heart disease  
 cardiovascular risk

Japan

lipid blood level

fatty acid blood level

lipid analysis

human

male

female

human experiment

normal human

aged

adult

article

Drug Descriptors:

\*linoleic acid

\*perilla oil

\*oxidized low density lipoprotein

\*lipid

\*fatty acid

palmitic acid

stearic acid

oleic acid

RN (linoleic acid) 1509-85-9, 2197-37-7, 60-33-3, 822-17-3; (lipid)  
 66455-18-3; (palmitic acid) 57-10-3; (stearic acid) 57-11-4, 646-29-7;  
 (oleic acid) 112-80-1, 115-06-0

L3 ANSWER 14 OF 22 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 2000063476 EMBASE

TI Homocysteine and oxidized low density lipoprotein enhance platelet  
 adhesion to endothelial cells under flow conditions: Distinct mechanisms  
 of thrombogenic modulation.

AU Dardik R.; Varon D.; Tamarin I.; Zivelin A.; Salomon O.; Shenkman B.;  
 Savion N.

CS Dr. R. Dardik, National Hemophilia Center, Sheba Medical Center, Tel  
 Hashomer 52621, Israel

SO Thrombosis and Haemostasis, (2000) 83/2 (338-344).

Refs: 30

ISSN: 0340-6245 CODEN: THHADQ

CY Germany

DT Journal; Article

FS 025 Hematology

037 Drug Literature Index

LA English

SL English

AB We investigated the effects of two well established risk factors for  
 cardiovascular disease, homocysteine and oxidized low density lipoprotein

**antibody.** The enhancing effect of homocysteine was also greatly reduced by inhibition of the EC .alpha.(v).beta.3 integrin, but was not affected by blockade of EC ICAM-1. On the other hand, **ox-LDL**-induced enhancement in platelet - EC adhesion was greatly inhibited by blocking ICAM-1 or .alpha.(v).beta.3, but remained unaffected by inhibition of TF activity. Preincubation of platelets with the glycoprotein IIb-IIIa (GPIIb-IIIa) antagonist Reo-Pro has virtually abolished the enhancing effect of both homocysteine and **ox-LDL**. Our results suggest that homocysteine and **ox-LDL** might increase endothelial thrombogenicity by distinct mechanisms: homocysteine - by inducing TF activity, and **ox-LDL** - by upregulating ICAM-1, both of which enhance GPIIb-IIIa/**fibrinogen** dependent platelet adhesion to EC. The .alpha.(v).beta.3 integrin, although not affected by EC stimulation, seems to play a crucial role in platelet-EC interaction regardless of the mechanism of EC perturbation.

CT Medical Descriptors:

\*blood flow  
\*endothelium cell  
\*thrombocyte adhesion  
\*thrombogenesis  
cardiovascular disease: ET, etiology  
risk factor  
shear rate  
thrombocyte activation  
thrombogenicity: ET, etiology  
human

controlled study

human cell

article

priority journal

Drug Descriptors:

\*homocysteine

\*oxidized low density lipoprotein

abciximab: DV, drug development

**fibrinogen receptor: EC, endogenous compound**

**fibrinogen: EC, endogenous compound**

fibrin: EC, endogenous compound

integrin: EC, endogenous compound

intercellular adhesion molecule 1: EC, endogenous compound

**polyclonal antibody**

thromboplastin: EC, endogenous compound

RN (homocysteine) 454-28-4, 6027-13-0; (abciximab) 143653-53-6; (**fibrinogen**) 9001-32-5; (fibrin) 9001-31-4; (intercellular adhesion molecule 1) 126547-89-5; (thromboplastin) 9035-58-9

CN Reopro

L3 ANSWER 15 OF 22 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 1999372263 EMBASE

TI Oxidized low-density lipoprotein inhibits the binding of monoclonal **antibody** to platelet glycoprotein IIB-IIIA.

AU Szuwart T.; Zhao B.; Fritsch A.; Mertens K.; Dierichs R.

CS Dr. R. Dierichs, Platelet Research Unit, Institute of Anatomy, Vesaliusweg 2-4, D-48149 Munster, Germany. dierich@uni-muenster.de

SO Thrombosis Research, (1999) 96/2 (85-90).

Refs: 31

ISSN: 0049-3848 CODEN: THBRAA

PUI S 0049-3848(99)00088-2

CY United Kingdom

DT Journal; Article

FS 018 Cardiovascular Diseases and Cardiovascular Surgery

025 Hematology

029 Clinical Biochemistry

030 Pharmacology

037 Drug Literature Index

005 General Pathology and Pathological Anatomy

LA English

SL English

AB Previous studies have shown that oxidized low-density lipoprotein (LDL) induces platelet activation more effectively than native LDL. To achieve a better understanding of the mechanism underlying the activation of human platelets by **oxidized LDL**, the present study relates the effect of oxidized LDL to changes of binding

\*atherogenesis: ET, etiology  
 ultrastructure  
 electrophoresis  
 drug distribution  
 human  
 controlled study  
 human cell  
 article  
 priority journal  
 Drug Descriptors:  
 \*oxidized low density lipoprotein: PD, pharmacology  
 \*oxidized low density lipoprotein: EC, endogenous compound  
 \*monoclonal antibody: PD, pharmacology  
 \*monoclonal antibody: PK, pharmacokinetics  
 \*fibrinogen receptor: PD, pharmacology

L3 ANSWER 16 OF 22 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

AN 1998079320 EMBASE

TI 3-Nitrotyrosine in the proteins of human plasma determined by an ELISA method.

AU Khan J.; Brennan D.M.; Bradley N.; Gao B.; Bruckdorfer R.; Jacobs M.

CS M. Jacobs, Department of Pharmacology, Royal Free Hospital, School of Medicine, Rowland Hill Street, London NW3 2PF, United Kingdom

SO Biochemical Journal, (1 Mar 1998) 330/2 (795-801).

Refs: 31

ISSN: 0264-6021 CODEN: BIJOAK

CY United Kingdom

DT Journal; Article

FS 029 Clinical Biochemistry

LA English

SL English

AB The modification of tyrosine residues in proteins to 3-nitrotyrosine by peroxynitrite or other potential nitrating agents has been detected in biological systems that are subject to oxidative stress. A convenient semi-quantitative method has been developed to assay nitrated proteins in biological fluids and homogenates using a competitive ELISA developed in our laboratory. This assay selectivity detected 3-nitro-L-tyrosine residues in a variety of peroxynitrite-treated proteins (BSA, human serum albumin (HSA), .alpha.1-antiprotease inhibitor, pepsinogen and **fibrinogen**) and also in a nitrated peptide, but had a low affinity for free 3-nitro-L-tyrosine and 3-chloro-L-tyrosine. The IC50 values for the inhibition of **antibody** binding by different nitrated proteins were in the range 5-100 nM, suggesting that the **antibody** discriminated between nitrotyrosine residues in different environments. The presence of nitrotyrosine in plasma proteins was detected by Western blot analysis and quantified by the ELISA. A concentration of 0.12 +/- 0.01 .mu.M nitro-BSA equivalents was measured in the proteins of normal plasma which was increased in peroxynitrite-treated plasma and was elevated in inflammatory conditions. HSA and low-density lipoprotein (LDL) isolated from plasma contained 0.085 +/- 0.04 and 0.03 +/- 0.006 nmol nitro-BSA equivalents/mg protein, respectively. Comparison of the level of nitration in peroxynitrite-treated HSA and LDL in the presence and absence of plasma indicates that nitration and presumably oxidation is inhibited by plasma antioxidants. The presence of nitrotyrosine in LDL is consistent with previous reports implicating peroxynitrite in the oxidative modification of lipoproteins and the presence of a low concentration of **oxidized LDL** in the blood.

CT Medical Descriptors:

\*enzyme linked immunosorbent assay  
 amino acid analysis  
 protein determination  
 antigen binding  
 immunoblotting  
 inflammation  
 lipoprotein blood level  
 nitration  
 protein blood level  
 human  
 normal human  
 adult  
 article  
 priority journal

TI Platelet integrin .alpha.(IIb).beta.3 (GPIIb-IIIa) is not implicated in  
the binding of LDL to intact resting platelets.  
AU Pedreno J.; Fernandez R.; Cullare C.; Barcelo A.; Elorza M.A.; De  
Castellarnau C.  
CS Dr. J. Pedreno, FIC (Pabellon Cardiologia), Hospital de la Santa Creu i  
Sant Pau, Avenida San Antonio Maria Claret 167, 08025 Barcelona, Spain  
SO Arteriosclerosis, Thrombosis, and Vascular Biology, (1997) 17/1 (156-163).

Refs: 47  
ISSN: 1079-5642 CODEN: ATVBFA

CY United States  
DT Journal; Article  
FS 029 Clinical Biochemistry  
LA English  
SL English

AB It has been suggested that the **fibrinogen** receptor (glycoprotein [GP] IIb-IIIa or platelet integrin .alpha.(IIb).beta.3) could be the binding site for low-density lipoprotein (LDL); however, recent data do not support this. Furthermore, GPIIb and not the GPIIb-IIIa complex is the main binding protein for lipoprotein(a) [Lp(a)]. In the present study, we have investigated the interaction between Lp(a) particles and platelet LDL binding sites and whether platelet integrin .alpha.(IIb).beta.3 is implicated. Displacement experiments showed that 125I-LDL binding to intact resting platelets was inhibited with the same apparent affinity by both unlabeled LDL and apolipoprotein(a)-free lipoprotein particles [Lp(a)], an LDL-like particle prepared from Lp(a)]. Hill coefficients for displacement curves suggested that a single set of binding sites was involved. In contrast, both native and oxidized Lp(a) particles were unable to inhibit platelet LDL binding. Furthermore, platelets bound 125I-Lp(a)-particles to a class of saturable binding sites numbering approximately 1958 +/- 235 binding sites per platelet with a dissociation constant (K(d) of 48.3 +/- 12 x 10<sup>-9</sup> mol/L. These values were similar to those obtained for LDL. In contrast to Lp(a), evidence indicates that platelet integrin .alpha.(IIb).beta.3 was not involved in the interaction of LDL and intact resting platelets. First, specific ligands for platelet integrin .alpha.(IIb).beta.3, such as **fibrinogen**, vitronectin, and fibronectin, were unable to inhibit the binding of LDL to intact resting platelets. Second, similar LDL binding characteristics (K(d) and B(max) values) were found in platelets from control subjects and patients with type I and type II Glanzmann's thrombasthenia, characterized by total and partial lack of GPIIb-IIIa and **fibrinogen**, respectively. Third, polyclonal **antibodies** against the GPIIb-IIIa complex (edu-3 and 5B12), human antisera against platelet alloantigens (anti-Bak(a/B) and anti-PL(A1/2), anti-integrin subunits (anti-.alpha.) (v) and anti-.beta.3, and a wide panel of monoclonal **antibodies** (mAbs) against well-known epitopes of GPIIb (M3, M4, M5, M6, and M95-2b) and GPIIIa (P23-7, P33, P37, P40 and P97) did not affect platelet LDL binding. Finally, in contrast to the proaggregatory effect of native and **oxidized LDL**, both native and oxidized Lp(a) particles caused a significant dose-dependent decrease of collagen-induced platelet aggregation. In conclusion, we demonstrate that neither the GPIIb-IIIa complex nor GPIIb and GPIIIa individually are membrane binding proteins for LDL on intact resting platelets. Lp(a) particles do not interact with platelet LDL binding sites, and their biological response is clearly different from that of LDL.

CT Medical Descriptors:  
\*cell adhesion  
\*thrombocyte  
article  
binding affinity  
binding site  
dissociation constant  
glanzmann disease  
human  
human cell  
ligand binding  
membrane binding  
priority journal  
thrombocyte aggregation  
Drug Descriptors:  
\***fibrinogen receptor**  
\*integrin  
\*lipoprotein a

SO FREE RADICAL BIOLOGY AND MEDICINE, (2001 Feb 1) 30 (3) 277-86.  
 Journal code: FRE; 8709159. ISSN: 0891-5849.  
 CY United States  
 DT Journal; Article; (JOURNAL ARTICLE)  
 LA English  
 FS Priority Journals  
 EM 200104  
 ED Entered STN: 20010502  
 Last Updated on STN: 20010502  
 Entered PubMed: 20010222  
 Entered Medline: 20010426  
 AB **Antibodies** against malondialdehyde (MDA)-modified proteins are often increased in patients with diseases related to oxidative stress. However, the clinical significance of these **antibodies** is hampered by their frequent presence also in healthy controls. Aim of this work has been to characterize the immune reactivity against MDA-derived antigens in healthy subjects. The sera of 120 healthy subjects contained IgG and IgM targeting MDA-modified human albumin (HSA), **fibrinogen**, and LDL. These sera also displayed weak reactivity with **oxidized LDL** and HSA complexed with oxidized arachidonic acid. Conversely, oxidized HSA or HSA complexed with other aldehydic lipid peroxidation products was not recognized. Control sera also did not recognize cyclic dihydropyridine-MDA products, while HSA-MDA reactivity was associated ( $r > 0.9$ ;  $p < .0005$ ) with the presence of fluorescent lysine-conjugated-imine cross-links. In Western blots both IgG and IgM recognized high molecular weight HSA-MDA aggregates, but not monomeric HSA-MDA adducts. The addition of sodium cyanoborohydride, that prevented conjugated-imine fluorescence and protein aggregation during HSA-MDA preparation, abolished the **antibody** binding. This suggested that the plasma of healthy subjects contained IgG and IgM recognizing protein aggregates linked through 1-amino-3-imino-propene bridges. The function of these **antibodies** is at the moment unknown, but they might contribute to scavenging MDA cross-linked proteins.  
 CT Check Tags: Female; Human; Male; Support, Non-U.S. Gov't  
 Adult  
 Arachidonic Acid: CH, chemistry  
 Arachidonic Acid: IM, immunology  
 \*Autoantibodies: BL, blood  
 Autoantigens: IM, immunology  
 \*Blood Proteins: CH, chemistry  
 \*Blood Proteins: IM, immunology  
 Blotting, Western  
**Fibrinogen: CH, chemistry**  
**Fibrinogen: IM, immunology**  
 IgG: BL, blood  
 IgM: BL, blood  
 Lipoproteins, LDL: CH, chemistry  
 Lipoproteins, LDL: IM, immunology  
 \*Malondialdehyde: CH, chemistry  
 \*Malondialdehyde: IM, immunology  
 Middle Age  
 Serum Albumin: CH, chemistry  
 Serum Albumin: IM, immunology  
 RN 506-32-1 (Arachidonic Acid); 542-78-9 (Malondialdehyde); **9001-32-5 (Fibrinogen)**  
 CN 0 (Autoantibodies); 0 (Autoantigens); 0 (Blood Proteins); 0 (IgG); 0 (IgM); 0 (Lipoproteins, LDL); 0 (Serum Albumin); 0 (oxidized low density lipoprotein)  
 L3 ANSWER 19 OF 22 MEDLINE  
 AN 2000202003 MEDLINE  
 DN 20202003 PubMed ID: 10739396  
 TI Homocysteine and oxidized low density lipoprotein enhanced platelet adhesion to endothelial cells under flow conditions: distinct mechanisms of thrombogenic modulation.  
 AU Dardik R; Varon D; Tamarin I; Zivelin A; Salomon O; Shenkman B; Savion N  
 CS National Hemophilla Center and Institute of Thrombosis and Hemostasis, Sheba Medical Center, Tel Hashomer, Israel.  
 SO THROMBOSIS AND HAEMOSTASIS, (2000 Feb) 83 (2) 338-44.  
 Journal code: VQ7; 7608063. ISSN: 0340-6245.  
 CY GERMANY: Germany, Federal Republic of  
 DT Journal; Article; (JOURNAL ARTICLE)

slightly reduced the expression of the intercellular adhesion molecule I (ICAM-1). In contrast, **ox-LDL** treatment upregulated ICAM-1 expression and had no significant effect on endothelial TF activity. Neither homocysteine nor **Ox-LDL** affected surface expression of the alpha(v)beta3 integrin. The homocysteine-induced enhancement in platelet adhesion was almost completely abolished by blockade of the EC TF activity by a polyclonal **antibody**. The enhancing effect of homocysteine was also greatly reduced by inhibition of the EC alpha(v)beta3 integrin, but was not affected by blockade of EC ICAM-1. On the other hand, **ox-LDL**-induced enhancement in platelet - EC adhesion was greatly inhibited by blocking ICAM-1 or alpha(v)beta3, but remained unaffected by inhibition of TF activity. Preincubation of platelets with the glycoprotein IIb-IIIa (GPIIb-IIIa) antagonist Reo-Pro has virtually abolished the enhancing effect of both homocysteine and **ox-LDL**. Our results suggest that homocysteine and **ox-LDL** might increase endothelial thrombogenicity by distinct mechanisms: homocysteine - by inducing TF activity, and **ox-LDL** - by upregulating ICAM-1, both of which enhance GPIIb-IIIa/**fibrinogen** dependent platelet adhesion to EC. The alpha(v)beta3 integrin, although not affected by EC stimulation, seems to play a crucial role in platelet-EC interaction regardless of the mechanism of EC perturbation.

CT Check Tags: Human; Support, Non-U.S. Gov't

**Antibodies: PD, pharmacology**

**Antibodies, Monoclonal: PD, pharmacology**

Calcium: PD, pharmacology

Cell Adhesion Molecules: BI, biosynthesis

Cell Adhesion Molecules: DE, drug effects

Endothelium, Vascular: CH, chemistry

Endothelium, Vascular: CY, cytology

\*Endothelium, Vascular: ME, metabolism

Fibrin: BI, biosynthesis

Fibrin: PH, physiology

**Fibrinogen: PD, pharmacology**

\*Homocysteine: PD, pharmacology

Homocysteine: PH, physiology

Immunoglobulins, Fab: PD, pharmacology

Intercellular Adhesion Molecule-1: BI, biosynthesis

Intercellular Adhesion Molecule-1: DE, drug effects

\*Lipoproteins, LDL: PD, pharmacology

Lipoproteins, LDL: PH, physiology

Oxidation-Reduction

\*Platelet Adhesiveness: DE, drug effects

Platelet Aggregation Inhibitors: PD, pharmacology

Platelet Glycoprotein GPIIb-IIIa Complex: AI, antagonists & inhibitors

Platelet Glycoprotein GPIIb-IIIa Complex: PD, pharmacology

Receptors, Cell Surface: BI, biosynthesis

Receptors, Cell Surface: DE, drug effects

Receptors, Vitronectin: BI, biosynthesis

Receptors, Vitronectin: DE, drug effects

Receptors, Vitronectin: ME, metabolism

Thromboplastin: BI, biosynthesis

Thromboplastin: DE, drug effects

Thromboplastin: IM, immunology

Umbilical Veins: CY, cytology

RN 126547-89-5 (Intercellular Adhesion Molecule-1); 143653-53-6 (abciximab);

454-28-4 (Homocysteine); 7440-70-2 (Calcium); 9001-31-4 (Fibrin);

**9001-32-5 (Fibrinogen)**; 9035-58-9 (Thromboplastin)

CN 0 (**Antibodies**); 0 (**Antibodies, Monoclonal**); 0 (Cell

Adhesion Molecules); 0 (Immunoglobulins, Fab); 0 (Lipoproteins, LDL); 0

(Platelet Aggregation Inhibitors); 0 (Platelet Glycoprotein GPIIb-IIIa

Complex); 0 (Receptors, Cell Surface); 0 (Receptors, Vitronectin); 0

(oxidized low density lipoprotein)

L3 ANSWER 20 OF 22 MEDLINE

AN 2000199490 MEDLINE

DN 20199490 PubMed ID: 10737229

TI Long-term effects of dietary alpha-linolenic acid from perilla oil on serum fatty acids composition and on the risk factors of coronary heart disease in Japanese elderly subjects.

AU Ezaki O; Takahashi M; Shigematsu T; Shimamura K; Kimura J; Ezaki H; Gotoh

T

CS Division of Clinical Nutrition, National Institute of Health and

ELISA using **antibody** specific to OxLDL. By merely replacing soybean cooking oil (SO) with perilla oil (PO) (i.e., increasing 3 g/d of ALA), the n-6/n-3 ratio in the diet was changed from 4:1 to 1:1. Twenty Japanese elderly subjects were initially given a SO diet for at least 6 mo (baseline period), a PO diet for 10 mo (intervention period), and then returned to the previous SO diet (washout period). ALA in the total serum lipid increased from 0.8 to 1.6% after 3 mo on the PO diet, but EPA and DHA increased in a later time, at 10 mo after the PO diet, from 2.5 to 3.6% and 5.3 to 6.4%, respectively ( $p < 0.05$ ), and then returned to baseline in the washout period. In spite of increases of serum n-3 fatty acids, the OxLDL concentration did not change significantly when given the PO diet. Body weight, total serum cholesterol, triacylglycerol, glucose, insulin and HbA1c concentrations, platelet count and aggregation function, prothrombin time, partial thromboplastin time, **fibrinogen** and PAI-1 concentration, and other routine blood analysis did not change significantly when given the PO diet. These data indicate that, even in elderly subjects, a 3 g/d increase of dietary ALA could increase serum EPA and DHA in 10 mo without any major adverse effects.

CT Check Tags: Female; Human; Male; Support, Non-U.S. Gov't

Aged

Aged, 80 and over

Cookery

\*Coronary Disease: ET, etiology

\*Dietary Fats: PD, pharmacology

Enzyme-Linked Immunosorbent Assay

\*Fatty Acids: BL, blood

Lipids: BL, blood

Lipoproteins: BL, blood

Lipoproteins, LDL: BL, blood

\*Plant Oils: PD, pharmacology

Risk Factors

\*alpha-Linolenic Acid: PD, pharmacology

RN 463-40-1 (alpha-Linolenic Acid); 68132-21-8 (perilla seed oil)

CN 0 (Dietary Fats); 0 (Fatty Acids); 0 (Lipids); 0 (Lipoproteins); 0

(Lipoproteins, LDL); 0 (Plant Oils); 0 (oxidized low density lipoprotein)

L3 ANSWER 21 OF 22 MEDLINE

AN 1998149755 MEDLINE

DN 98149755 PubMed ID: 9480893

TI 3-Nitrotyrosine in the proteins of human plasma determined by an ELISA method.

CM Erratum in: Biochem J 1998 Jun 15;332 (Pt 3):808

Erratum in: Brennan DM[corrected to Brennand DM]

AU Khan J; Brennand D M; Bradley N; Gao B; Bruckdorfer R; Jacobs M; Brennan D M

CS Department of Pharmacology, Royal Free Hospital School of Medicine, Rowland Hill Street, London NW3 2PF, U.K.

SO BIOCHEMICAL JOURNAL, (1998 Mar 1) 330 ( Pt 2) 795-801.

Journal code: 9YO; 2984726R. ISSN: 0264-6021.

CY ENGLAND: United Kingdom

DT Journal; Article; (JOURNAL ARTICLE)

LA English

FS Priority Journals

EM 199804

ED Entered STN: 19980422

Last Updated on STN: 19990129

Entered Medline: 19980416

AB The modification of tyrosine residues in proteins to 3-nitrotyrosine by peroxy-nitrite or other potential nitrating agents has been detected in biological systems that are subject to oxidative stress. A convenient semi-quantitative method has been developed to assay nitrated proteins in biological fluids and homogenates using a competitive ELISA developed in our laboratory. This assay selectivity detected 3-nitro-L-tyrosine residues in a variety of peroxy-nitrite-treated proteins (BSA, human serum albumin (HSA), alpha1-antiprotease inhibitor, pepsinogen and **fibrinogen**) and also in a nitrated peptide, but had a low affinity for free 3-nitro-L-tyrosine and 3-chloro-L-tyrosine. The IC50 values for the inhibition of **antibody** binding by different nitrated proteins were in the range 5-100 nM, suggesting that the **antibody** discriminated between nitrotyrosine residues in different environments.

The presence of nitrotyrosine in plasma proteins was detected by Western blot analysis and quantified by the ELISA. A concentration of 0.12 +/- 0.01 microM nitro-BSA equivalents was measured in the proteins of normal plasma



Serum Albumin: CH, chemistry  
 \*Tyrosine: AA, analogs & derivatives  
 Tyrosine: AN, analysis  
 RN 3604-79-3 (3-nitrotyrosine); 55520-40-6 (Tyrosine)  
 CN 0 (Blood Proteins); 0 (Lipoproteins, LDL); 0 (Serum Albumin)

L3 ANSWER 22 OF 22 MEDLINE  
 AN 97164889 MEDLINE  
 DN 97164889 PubMed ID: 9012651  
 TI Platelet integrin alpha IIb beta 3 (GPIIb-IIIa) is not implicated in the binding of LDL to intact resting platelets.  
 AU Pedreno J; Fernandez R; Cullare C; Barcelo A; Elorza M A; de Castellarnau C  
 CS Department of Biochemistry, Hospital Universitario Son Dureta, Palma de Mallorca, Spain.  
 SO ARTERIOSCLEROSIS, THROMBOSIS, AND VASCULAR BIOLOGY, (1997 Jan) 17 (1) 156-63.  
 Journal code: B89; 9505803. ISSN: 1079-5642.  
 CY United States  
 DT Journal; Article; (JOURNAL ARTICLE)  
 LA English  
 FS Priority Journals  
 EM 199702  
 ED Entered STN: 19970306  
 Last Updated on STN: 19970306  
 Entered Medline: 19970225

AB It has been suggested that the **fibrinogen** receptor (glycoprotein [GP] IIb-IIIa or platelet integrin alpha IIb beta 3) could be the binding site for low-density lipoprotein (LDL); however, recent data do not support this. Furthermore, GPIIb and not the GPIIb-IIIa complex is the main binding protein for lipoprotein(a) [Lp(a)]. In the present study, we have investigated the interaction between Lp(a) particles and platelet LDL binding sites and whether platelet integrin alpha IIb beta 3 is implicated. Displacement experiments showed that 125I-LDL binding to intact resting platelets was inhibited with the same apparent affinity by both unlabeled LDL and apolipoprotein(a)-free lipoprotein particles [Lp(a)-, an LDL-like particle prepared from Lp(a)]. Hill coefficients for displacement curves suggested that a single set of binding sites was involved. In contrast, both native and oxidized Lp(a) particles were unable to inhibit platelet LDL binding. Furthermore, platelets bound 125I-Lp(a)- particles to a class of saturable binding sites numbering approximately 1958 +/- 235 binding sites per platelet with a dissociation constant (Kd) of 48.3 +/- 12 x 10<sup>-9</sup> mol/L. These values were similar to those obtained for LDL. In contrast to Lp(a), evidence indicates that platelet integrin alpha IIb beta 3 was not involved in the interaction of LDL and intact resting platelets. First, specific ligands for platelet integrin alpha IIb beta 3, such as **fibrinogen**, vitronectin, and fibronectin, were unable to inhibit the binding of LDL to intact resting platelets. Second, similar LDL binding characteristics (Kd and Bmax values) were found in platelets from control subjects and patients with type I and type II Glanzmann's thrombasthenia, characterized by total and partial lack of GPIIb-IIIa and **fibrinogen**, respectively. Third, polyclonal **antibodies** against the GPIIb-IIIa complex (edu-3 and 5B12), human antiserums against platelet alloantigens (anti-Baka/B and anti-PLA1/2), anti-integrin subunits (anti-alpha v and anti-beta 3), and a wide panel of monoclonal **antibodies** (mAbs) against well-known epitopes of GPIIb (M3, M4, M5, M6, and M95-2b) and GPIIIa (P23-7, P33, P37, P40, and P97) did not affect platelet LDL binding. Finally, in contrast to the proaggregatory effect of native and **oxidized LDL**, both native and oxidized Lp(a) particles caused a significant dose-dependent decrease of collagen-induced platelet aggregation. In conclusion, we demonstrate that neither the GPIIb-IIIa complex nor GPIIb and GPIIIa individually are membrane binding proteins for LDL on intact resting platelets. Lp(a) particles do not interact with platelet LDL binding sites, and their biological response is clearly different from that of LDL.

CT Check Tags: Human; Support, Non-U.S. Gov't  
 Binding Sites  
 \*Blood Platelets: ME, metabolism  
 \*Lipoproteins, LDL: ME, metabolism  
 \*Platelet Glycoprotein GPIIb-IIIa Complex: ME, metabolism  
 Radioligand Assay

CN 0 (Lipoproteins, LDL); 0 (Platelet Glycoprotein GPIIb-IIIa Complex)